Inter-Simple Sequence Repeat-Restriction Fragment Length Polymorphisms for DNA Fingerprinting

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As part of this laboratory’s honeybee genetics and breeding program, DNA markers are needed for uses such as the identification of populations and species and marker-assisted breeding. In particular, DNA markers are needed to discriminate *Varroa*-mite-resistant Russian honey bees, imported from far-eastern Russia, and their offspring (1), from the other honey bees found in the USA. I tried RAPDs, but they were often unrepeatable. AFLPs require an undesirably large amount of preparation effort for each sample. Microsatellites have been useful for other studies in honey bees (2,3), but for this purpose there is inadequate variation with the primer pairs available. Microsatellite-primed PCR involves the amplification of DNA using a single primer composed of a microsatellite sequence (simple sequence repeat; SSR) with 1–3 selective, often degenerate, nucleotides at the end. The amplified fragments are termed inter-simple sequence repeats (4). The first report of ISSRs by Zietkiewicz et al. (5) discussed several mammals, other vertebrates, plants, and *E. coli*. Since then, ISSRs have been reported mostly from studies in plants, particularly where inadequate variation or reproducibility were available with other methods [e.g., primer pairs in *Ipomoea* (4) and in *Fragaria* linkage analysis (6), cocoa germplasm characterization (7), inter-simple sequence repeat (ISSR) inheritance in citrus (8), and classifying rice germplasm (9)]. However, ISSRs have also been used to study variation in aphids, a mosquito, and a rotifer (10); a rice pest insect (11); the silkworm (12); and corals (13). I also successfully amplified ISSR fragments from *Varroa* mites and Hawaiian *Drosophila* (data not reported). This indicates that ISSRs and ISSR-RFLPs should be detectable in any organism where DNA markers can be detected. Here I describe a new method to increase the amount of detectable DNA variation by digesting ISSRs with restriction enzymes to produce ISSR-RFLPs. The digested amplification products are detected using ethidium bromide after electrophoresis. This is a novel combination of procedures and is an improvement over just ISSRs because restriction enzyme digestion of ISSR fragments allows the detection of significant amounts of additional variation using a more repeatable and reproducible procedure. This would be particularly useful in cases where inadequate amounts of detectable genetic variation are available using ISSRs or other methods or where more variation is desired from a limited number of ISSR primers. Detection of RFLPs in PCR-amplified DNA using ethidium bromide eliminates the requirements for Southern transfers, unique probes for detection, and labeled probes or primers. This simplifies detection and reduces costs.

Genomic DNA from the thorax of a single worker honey bee, *Apis mellifera* L., from a colony, with each colony from a different source, was extracted with 400 µL 10% Chelex® resin (14). SSR primers were purchased from the Oligonucleotide Synthesis Laboratory, NAPS Unit, University of British Columbia (Vancouver, BC, Canada). For initial evaluations of primers and DNA templates, the PCR procedure of Huang and Sun (4) was followed, but unlabeled primers were used and PCR products were detected by ethidium bromide staining after electrophoresis. Amplification was carried out in a total volume of 5 µL containing 0.35 µL Chelex extract template DNA, 0.21 µM primer, 0.2 mM dNTPs, 1.5 mM MgCl₂, 0.5 U Taq DNA polymerase...
PCR was performed in a PTC-100™ thermal cycler (MJ Research, Waltham, MA, USA) with a heated lid, using the following cycle profile: 1 cycle at 94°C for 4 min, followed by 34 cycles of 94°C for 45 s, 51°C for 45 s, 72°C for 1.5 min, and a final 7-min extension at 72°C. Primers often required individual optimization for concentrations of primer, dNTP, and Taq DNA polymerase. Following PCR, each amplification, along with 2 µL Elchrom loading buffer, was loaded into a precast 6% Poly(NAT)® horizontal slab gel (similar to acrylamide) in 30 mM TAE in an Elchrom SEA 2000® electrophoresis unit (Elchrom, Cham, Switzerland) and electrophoresed for 1 h at 105 V. For data recording, each gel was removed from the plastic backing, agitated for 40 min in 0.5 µg/mL ethidium bromide solution, and photographed on a UV transilluminator with a Kodak® EDAS 290 (Eastman Kodak, Rochester, NY, USA). For restriction enzyme digestion, PCR amplification was carried out in a total volume of 75 µL at the same reagent ratios as the 5-µL reactions. A 4-µL aliquot from each amplification was analyzed to see if the PCR amplifications were successful. If successful, 5-µL aliquots from the same amplification for each honey bee were digested with restriction enzymes (RE) (New England Biolabs, Beverly, MA, USA) as follows: 5 µL PCR amplification, 1 µg BSA if required, 1 µL 10× RE buffer, 2.5 U RE, water to 10 µL total. Restriction enzyme digestion was for 2 h or overnight at the temperature specified by the manufacturer, generally 37°C. Two microliters of Elchrom loading buffer were added to each digestion, and they were gel analyzed as above, except that electrophoresis was carried out for 50 min to avoid running small fragments off the gel.

Figure 1 shows the ISSR banding patterns produced by UBC primer 827, (AC)₈G, with three Russian (R) and three Italian-ancestry (nR) honey bees, all from different colonies. Figure 2 shows the ISSR-RFLP bands from restriction enzyme digestion of the ISSR bands in lanes 1 and 2. The odd-numbered lanes of Figure 2 are aliquots from the same PCR amplification as lane 3 of Figure 1. The even-numbered lanes of Figure 2 are aliquots from the same PCR amplification as lane 4 of Figure 1. Therefore, all restriction fragments are from digestion of the ISSR fragments from the two PCR amplifications shown in lanes 1 and 2, and variations are not nonspecific bands or artifacts of different amplifications. While some variation among the samples is present in Figure 1, lanes 3 and 4 differ
only by the presence of two additional lighter bands in lane 4. When the ISSR bands from these two samples are digested with 11 restriction enzymes, further variation is detectable between the pairs for all restriction enzymes (Figure 2). Thus, restriction enzyme digestion of these ISSR bands detects much additional DNA variation specific to these two bees without further PCR optimization for new primers.

With ethidium bromide detection, 2–6 prominent ISSR fragments and several fainter fragments were usually found when many different ISSR primers were tested (data not reported). Most of the fragments were in the 400–2000 bp range compared to the 50–500 bp range reported by Huang and Sun (4). For most primers, the patterns of the prominent fragments were much more repeatable at varying levels of template DNA than I had found with RAPDs. Several RFLPs were observed for most primer/restriction enzyme combinations. Thus restriction enzyme digestion detects much additional variation among ISSR PCR fragments.

Much of the initial effort in PCR-based studies involves optimization of the PCR conditions, and this also applies to ISSRs. Restriction enzyme digestion of ISSR bands does not require optimization for different ISSRs. Thus, much less optimization effort is required to screen additional restriction enzymes for one ISSR than is required to screen additional ISSR primers to detect an equal amount of variation. While the initial step to detect ISSR-RFLPs does require large reaction PCR to amplify the ISSR bands, PCR machines are not needed for restriction enzyme digestion. Thus, much less use of PCR machines is required for ISSR-RFLPs than for detection of an equal amount of ISSR variation. Effort can then be concentrated on those ISSR primers that are the most reliable and produce the most useful bands. While PCR amplifications with some ISSR primers often fail or are of uneven quality, restriction enzyme digestions of ISSR amplifications are much more reliable and repeatable.

Verification of successful PCR amplification before restriction enzyme digestion allows the researcher to avoid attempting to digest poor or failed amplifications. Failed or poor amplifications can be repeated or modified and this also applies to different samples can be substituted, minimizing waste of resources by identifying problems early in the process. This uncouples the relatively reliable process of restriction enzyme digestion from the often more problematic process of PCR, thus increasing the reliability of the final data.

For research and stock identification purposes, we needed DNA markers to distinguish Russian honey bees from other types of honey bees present in the USA. Using a different primer and four restriction enzymes, it is possible to use ISSR-RFLPs to discriminate between Russian and other honey bees (data not reported).

REFERENCES

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